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Growth Phase-Dependent Response of *Helicobacter pylori* to Iron Starvation

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Iron is an essential nutrient that is often found in extremely limited available quantities within eukaryotic hosts. Because of this, many pathogenic bacteria have developed regulated networks of genes important for iron uptake and storage. In addition, it has been shown that many bacteria use available iron concentrations as a signal to regulate virulence gene expression. We have utilized DNA microarray technology to identify genes of the human pathogen *Helicobacter pylori* that are differentially regulated on a growth-inhibiting shift to iron starvation conditions. In addition, the growth phase-dependent expression of these genes was investigated by examining both exponential and stationary growth phase cultures. We identified known iron-regulated genes, as well as a number of genes whose regulation by iron concentration was not previously appreciated. Included in the list of regulated factors were the known virulence genes cagA, vacA, and napA. We examined the effect of iron starvation on the motility of H. pylori and found that exponential- and stationary-phase cultures responded differently to the stress. We further found that while growing cells are rapidly killed by iron starvation, stationary-phase cells show a remarkable ability to survive iron depletion. Finally, bioinformatic analysis of the predicted promoter regions of the differentially regulated genes led to identification of several putative Fur boxes, suggesting a direct role for Fur in iron-dependent regulation of these genes.

The essential nature of iron for virtually all organisms is a double-edged sword that requires a careful balancing act to ensure survival. On the one hand, iron is a required cofactor of many enzymes involved in the maintenance of basic metabolic functions such as electron transport. On the other hand, interaction of free oxygen with iron leads to the Fenton reaction, which results in the production of oxygen radicals that can cause severe damage to the majority of cellular biomolecules (65). Due to these two opposing phenomena, it is perhaps no surprise that organisms have developed tightly regulated systems for both uptake and sequestration of iron. Within the context of mammalian systems this is accomplished in part through the action of iron binding molecules such as transferrin, heme and lactoferrin (42, 44). In addition to protecting cells from the harmful effects of oxygen radicals, the ability to sequester iron also limits bacterial growth within the mammalian host. This is evidenced by extensive epidemiological data suggesting that there is a strong link between the iron status of the host and susceptibility to a number of bacterial pathogens

In light of the requirement for iron and the stiff competition for this nutrient, it is no surprise that successful pathogenic microorganisms have developed intricate systems that allow them to acquire and store iron within the bacterial cell. In fact, genes that encode factors involved in these processes are often considered to be virulence factors and have been the subject of intense study by numerous groups (9). Intrinsic to the ability to acquire iron is the fact that microbes must also be able to sense and respond to changing iron concentrations within the environment. This is crucial to help maintain the homeostatic relationship between having enough iron to grow and avoiding iron toxicity. In a number of organisms, this regulation is accomplished by the ferric uptake regulator (Fur) protein (15). Found in both gram-negative and gram-positive bacteria, Fur functions by binding to promoter regulatory elements, called Fur boxes, in an iron-dependent manner. Typically, Fur-regulated promoters are repressed under iron-replete conditions and derepressed under iron-depleted conditions. A notable exception to this simple mode of regulation has recently been shown in Helicobacter pylori. The H. pylori Fur protein controls the transcription of both iron uptake and storage genes, frpB and pfr, respectively (18). This is accomplished by alteration of binding specificity based on the iron status of the Fur protein.

Other factors of *H. pylori* are known to show iron-dependent regulation, although thorough characterization of the roles of many of these is lacking. In addition to binding to the *pfr* and *frpB* promoters, Fur directly binds and autoregulates its own promoter (17). Expression of the genes *fecA1*, *fecA2*, *feoB*, and *frpB2/3* is iron regulated in a Fur-dependent manner (16, 67), although it is unclear whether this dependency is direct or indirect. While all of these genes are suggested to play a role in iron uptake, a definitive role has been demonstrated only for *feoB* in *H. pylori* (68). Expression of a number of other proteins has also been shown to be regulated by iron. Among these, several outer membrane proteins (29, 31, 73), the vacuolating cytotoxin (*vacA*) (61), the riboflavin biosynthetic genes *ribAB*

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(72), and the fumarate reductase genes *frdCAB* have all been shown to exhibit changes in expression in response to varying concentrations of iron (20). Increased expression of the *vacA* toxin is in keeping with the finding that in addition to regulating genes that are important for iron uptake and homeostasis, many pathogenic organisms regulate virulence factors in response to iron concentration (9).

A thorough understanding of the genes involved in iron uptake and homeostasis is potentially of profound clinical importance because H. pylori colonization is associated with multiple forms of gastric disease. These range in severity from mild gastritis to duodenal and gastric ulcers and two forms of gastric malignancy, mucosa-associated lymphoid tissue lymphoma and adenocarcinoma (7, 22, 46). Additionally, H. pylori infection has been linked to iron deficiency anemia (IDA), particularly in children and premenopausal women (5, 53). Clinical evidence suggests that there are multiple mechanisms whereby H. pylori infection leads to IDA, either through the induction of impaired iron absorption by the host or through competition for the limited iron stores that are present (4). Even with approximately 50% of the world's population infected with H. pylori (37), the process of infection and survival of the bacterium within its gastric niche remains relatively poorly understood.

In a recent systematic global analysis of growth phase-dependent gene expression of H. pylori, the transcriptome of the bacterium was determined across the entire growth phase and revealed a dramatic switch in expression of many genes as bacteria made the transition from the exponential to the stationary phase of growth (63). Two major classes of genes affected at this transition were those known to encode virulence factors and those involved in iron homeostasis, suggesting the possibility that intracellular concentrations of iron may play a strong role in the modulation of both classes of genes. To better understand growth phase-modulated adaptive mechanisms utilized by H. pylori, we used spotted DNA microarrays to characterize the global response to iron starvation of both exponential- and stationary-phase cultures. Comparison of the transcription levels after growth-limiting iron chelation with those of the starting culture revealed three classes of genes: those affected specifically in the exponential phase or the stationary phase and those affected in both growth phases. Among these genes are many encoding proteins whose expression had previously been shown to be affected by iron concentration, as well as many whose iron-regulated gene expression had not previously been identified. Microscopic analysis of the effect of iron limitation on motility revealed that stationaryphase H. pylori cells show a marked ability to swim and survive for longer periods than do exponential-phase bacteria. Finally, analysis of the predicted promoter regions of the identified regulated genes led to the detection of several putative Fur boxes, thus suggesting a direct role for Fur in iron-dependent regulation of these genes.

MATERIALS AND METHODS

Bacterial strains and growth. $H.\ pylori$ strain SS1 (34) was maintained as a frozen stock at -80° C in brain heart infusion medium supplemented with 20% glycerol and 10% fetal bovine serum (FBS) (Gibco/BRL). Bacteria were grown on horse blood agar (HBA) plates containing 4% Columbia agar base (Oxoid), 5% defibrinated horse blood (HemoStat Labs, Dixon, Calif.), 0.2% B-cyclodextrin (Sigma), $10\ \mu g$ of vancomycin (Sigma) per ml, $5\ \mu g$ of cefsulodin (Sigma) per

ml, 2.5 U of polymyxin B (Sigma) per ml, 50 µg of cycloheximide (Sigma) per ml, 5 µg of trimethoprim (Sigma) per ml, and 8 µg of amphotericin B (Sigma) per ml under microaerophilic conditions at 37°C. A microaerobic atmosphere was generated using a CampyGen sachet (Oxoid) in a gas pack jar. For liquid culture, H. pylori was grown in brucella broth (BB) (Difco) containing 10% FBS with shaking in a microaerobic environment. The MIC of 2,2'-dipyridyl (DPP) that inhibited bacterial growth but not bacterial viability for the duration of the time courses was determined by supplementation of exponential-phase H. pylori cultures with increasing concentrations of DPP and subsequent monitoring of changes in the optical density at 600 nm OD600 and viable CFU by plating on HBA plates.

Iron chelation and RNA isolation. An overnight liquid culture of SS1 (OD₆₀₀ of 0.5 to 1.0) was harvested by centrifugation and resuspended to an OD_{600} of 0.05 to 0.1 in BB supplemented with 10% FBS. Bacterial samples were collected after dilution and represent the start time point. The master culture was then split to yield either two independent 70-ml cultures (chelation experiments) or four independent 70-ml cultures (supplementation experiments). The cultures were maintained under microaerobic conditions with shaking overnight at 37°C. Next, DPP was added to a concentration of 200 μM for each exponential-phase culture (OD $_{600}$ of 0.30 to 0.35) and stationary-phase culture (OD $_{600}$ of 0.80 to 0.90). Bacterial samples were collected immediately prior to the addition of DPP and represent the t = 0 time point. All cultures were maintained under microaerobic conditions with shaking at 37°C, and samples were harvested 15, 30, 60, and 90 min (experiment 1) or 30, 60, and 120 min (experiment 2) after iron chelation. At each time point, aliquots were removed from the medium, plated to determine CFU, examined microscopically to determine the level of motility and harvested on a 0.45-µm-pore-size cellulose filter by vacuum filtration. Filters containing the H. pylori cells were immediately frozen in liquid nitrogen and subsequently used for isolation of bacterial RNA. H. pylori RNA was isolated using TRIzol reagent (Gibco/BRL) as previously described (39). The RNA concentration was quantitated by determination of the OD260, and RNA integrity was verified by visualization on a 1% agarose gel.

Iron add-back. Growth of SS1 under iron-restricted conditions was conducted by first determining the concentration of DPP that would slow but not completely inhibit bacterial growth; 75 μ M was determined to be a suitable concentration. An overnight liquid culture of SS1 was used to inoculate BB supplemented with 10% FBS and 75 μ M DPP at a starting OD $_{600}$ of 0.05. The cultures were maintained under microaerobic conditions with shaking overnight at 37°C. After approximately 19 h, the culture had reached an OD $_{600}$ of 0.13. Iron was replenished in this culture by the addition of FeCl $_3$ to a final concentration of 1 mM, a concentration that was shown in an independent experiment to relieve iron-dependent growth restriction. The cultures were maintained under microaerobic conditions with shaking at 37°C. Samples were taken immediately prior to the addition of FeCl $_3$ (t=0 time point) and 30, 60, and 120 min thereafter, as described above, and subsequently used for RNA isolation and microarray analysis as described below.

Microarray hybridization and analysis. Equal concentration of each test RNA (t = 0, 15, 30, 60, 90, and 120 min from biologically independent experiments)and the reference RNA (t = 0 min) were used for cDNA synthesis in a standard reverse transcription reaction using Superscript II (Invitrogen) and Panorama H. pylori cDNA labeling primers (SigmaGenosys). Synthesized cDNA was purified using Qia-Quick PCR purification columns (Qiagen) as specified by the manufacturer and subsequently indirectly labeled with Cy5 (red) and Cy3 (green) fluorophores, respectively, as previously described (63). Individual Cy5 and Cy3 reactions were combined, and unincorporated dye was removed using a Qia-Quick PCR column as specified by the manufacturer. The eluates from the columns were concentrated by evaporation in a Speed Vac and resuspended in 11 μl of Tris-EDTA (TE). Then 1 μl of 25-mg/ml yeast tRNA, 2.55 μl of 20× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate), and 0.45 μl of 10% sodium dodecyl sulfate were added to each of the labeled probes. These were heated to 99°C for 2 min, cooled briefly, and then added to the H. pylori microarray for hybridization and stringency washes as previously described (47). The hybridized slides were scanned and analyzed using 4000A scanner and the GENEPIX 3.0 software (both from Axon).

Data were collated using the Stanford Microarray Database (54). For the chelation experiments, spots showing obvious abnormalities were excluded from analysis. Spot quality was further filtered by requiring that the Cy3 mean spot intensity/Cy5 median background intensity ratio be greater than or equal to 1.5 and the Cy5 mean spot intensity/Cy3 median background intensity ratio be greater than or equal to 1.5. Genes were filtered by requiring that the \log_2 of the red/green normalized ratio (mean) have an absolute value greater than or equal to 1 in at least one array for each experimental set. Biologically independent experiments were analyzed separately, and the data were collapsed to reveal

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factors whose expression was consistently altered by iron starvation. For the iron add-back experiment, spot quality was filtered by requiring that the Cy3 net spot intensity be greater than or equal to 100. Genes were filtered by requiring that the log₂ of the red/green normalized ratio (mean) have an absolute value greater than or equal to 0.59 in at least one array for each experimental set. Biologically independent experiments were analyzed separately, and the data were collapsed to reveal factors whose expression was consistently altered by iron addition. The list of factors was then cross-compared to the list generated for the iron chelation experiments, and the factors which overlap were obtained. All microarray data generated by this study are publicly available at http://genome-www.stanford.edu/microarray/.

Correlation coefficient determination. To verify the growth status of cultures used in the chelation experiments, arrays where the t=0 time point of the cultures was labeled in Cy5 and the start time point of the cultures was labeled in Cy3 for both the exponential- and stationary-phase time courses were compared to a previously published growth phase-dependent transcriptional profile of $H.\ pylori$ (63). Correlation coefficients were determined by comparing the \log_2 red/green ratios from these two arrays with the data from each of the arrays in the described control time course (63) by using Microsoft Excel.

Motility analysis and supplementation. For the motility and supplementation experiment, exponential- and stationary-phase cultures were grown and chelated as described above and maintained for extended periods. The motility of samples was monitored by live phase-contrast microscopy with glass slides and coverslips prewarmed to 37°C. A Hammamatsu C2400 video charge-coupled device camera was used to record movement in the field of view via an Argus-20 image processor (using the TRACE function) onto S-VHS video. Movement was traced over a 5-s period. Two sets of selected video frames for each time point were digitized for the generation of time-lapse films with the National Institutes of Health ObjectImage program. The percentage of motile bacteria at each time point was estimated using these films. The average percent motility data for each time point was plotted over time using Microsoft Excel.

Supplementation was performed at the 4-h time point postchelation and every 4 h thereafter by the removal of 5% of the culture and replacement with BB–FBS that had been chelated with 200 μ M DPP. All cultures were maintained under microaerobic conditions with shaking at 37°C. CFU were determined by plating on HBA plates.

Identification of predicted Fur binding sites. Each of the differentially regulated genes from the chelation experiment was visually examined to identify transcriptional organization in relation to the surrounding genes, and a list was generated that represented the predicted first gene in each putative operon identified. A Perl script was then written to extract the sequence of the 150 nucleotides immediately upstream of the predicted translational start site of the first gene in each predicted operon. A candidate Fur binding consensus sequence (AATAATNNTNA) for *H. pylori* was generated by aligning the high-affinity Fur binding sites of the *frpB* box (16) and *fur* box I (17) promoters, as well as the FURTA-Hp positive *fecA2* promoter (26), using Clustal X (62). The 150-nucleotide regions were then searched for this consensus motif by using an additional Perl script, and sequences showing 100% identity were aligned using Clustal X.

RESULTS AND DISCUSSION

Microarray analysis of iron-regulated gene expression. Determining a suitable DPP concentration that would not inhibit bacterial survival was a critical experimental parameter since decreased viability at later time points could greatly affect the transcriptional profiles obtained. Consequently, growth and survival curves were determined in BB supplemented with 10% FBS which had been chelated with increasing concentrations of DPP. The concentration determined to be suitable was $200~\mu\mathrm{M}$ as it inhibited H.~pylori growth and showed no effect on bacterial viability for time points up to several hours after chelation (data not shown).

Since previous microarray analysis had revealed that many of the genes that are subject to growth phase-dependent expression are known or predicted to play roles in iron uptake and homeostasis (63), we wished to assess the growth phase dependency of transcriptional changes induced by iron starvation. Since these previously observed differences were most

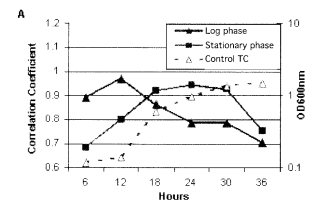
evident between the exponential and stationary phases, we depleted iron from cultures of *H. pylori* in these two growth phases as described in Materials and Methods. Bacterial cells were harvested during a time course up to 2 h postchelation, and total bacterial RNA was prepared and subsequently used for microarray analysis. To ensure reproducibility, chelation experiments were replicated in a separate experiment and only the results for genes that consistently showed at least a twofold change in expression in both experiments are reported.

To ensure that the described iron-chelated cultures were in fact in the exponential and stationary phases, respectively, the genome-wide transcriptional profile of each of the starting cultures was compared to the previously published growth phase-dependent transcriptome of *H. pylori* (63). In this previous work, the complete transcriptional profile of H. pylori as it transits the growth curve from early exponential phase to late stationary phase was determined. Using this information, we were able to calculate a genome-wide correlation coefficient between the t = 0 time points of our exponential and stationary iron chelation cultures and each of the time points during the growth curve. These results are plotted in Fig. 1A. This plot shows that the predicted exponential-phase t = 0 iron chelation array correlates most highly with the 12-h time point of the growth curve, which represents the early exponential phase. On the other hand, the predicted stationary-phase t = 0 iron chelation array correlates most highly with the 24-h time point of the growth curve, which represents the early stationary phase. This result was also confirmed by performing an unsupervised hierarchical cluster of the iron chelation data sets with the growth curve data. The dendrogram depicted in Fig. 1B clearly shows that in each case, the exponential- and stationaryphase iron chelation arrays cluster with their respective counterparts across the growth curve. These data indicate that the respective chelation cultures were in the desired growth phase for determination of the growth phase-dependent response to iron starvation.

Classification of iron-regulated genes. A total of 138 genes were shown to be differentially regulated by a minimum of twofold in the exponential-phase cultures (Table 1). The majority of these, 87 genes, were induced, while 51 genes showed a decrease in expression level. The transcriptional response of the stationary-phase cultures was much less dramatic, since only 75 genes displayed altered expression (Table 2). Of these, only 6 genes were repressed and the remaining 69 were induced. Direct comparison of the two tables revealed that only 30 genes overlapped between the exponential- and stationary-phase cultures (indicated by asterisks in Tables 1 and 2). Taken together, this indicates that a total of 183 individual genes showed altered expression on iron starvation of the two *H. pylori* cultures (Fig. 2).

Sorting of these genes into functional classes as defined by The Institute for Genomic Research revealed that a number of different biological functions are affected by the loss of iron. These include a large number of genes predicted to encode hypothetical or conserved proteins of unknown function, proteins involved in metabolism, components of the cell envelope or surface, and factors involved in transport and binding. For the sake of brevity, a few key genes are discussed below.

Iron starvation-repressed genes. (i) Iron sequestration factors. The *pfr* locus was found to be strongly repressed in the



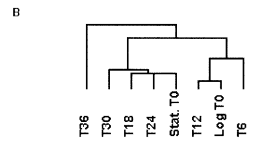


FIG. 1. (A) Verification of the growth phase of the exponentialand stationary-phase H. pylori cultures used in the chelation experiment. Representative correlation coefficients for the t = 0 time point of the iron chelation cultures were calculated by comparison with published expression profiles of H. pylori across a standard growth curve time course (Control TC) (63). The OD₆₀₀ of the Control TC is labeled on the right axis. Individual correlation coefficients between the exponential- and stationary-phase array data and the data for each of the arrays of the Control TC are indicated on the left-hand y axis. Time is indicated in hours on the x axis. The highest correlation coefficients for the indicated growth phase occur at t = 12 h and t = 1224 h and represent the point at which the iron chelation experiments were initiated. (B) Dendrogram showing segregation of exponentialphase (denoted Log T0) and stationary-phase (denoted Stat. T0) t = 0array data sets into respective branches generated by an unsupervised hierarchical cluster of the two t = 0 arrays from the chelation experiment with the array data from each of the time points of the Control TC. In each case, the chelation culture arrays cluster tightly with the concomitant growth phase arrays from the Control TC.

iron-starved stationary-phase cultures of *H. pylori*, validating the ability of our experimental design to identify known iron-regulated genes. Pfr acts as the major *H. pylori* ferritin and is responsible for the storage of iron in subcellular granules within the bacterial cytoplasm (6). The activity of this protein plays an important role in the *H. pylori* life cycle because it is required for growth and survival under iron starvation conditions. Additionally, Pfr activity is important for maximal iron uptake, since mutation of *pfr* greatly decreases the rate of transport of ⁵⁵FeCl₂ and ⁵⁵Fe³⁺ into the *H. pylori* cell (69). These activities are probably crucial within the tumultuous environment of the human stomach, since mutations in *pfr* result in an *H. pylori* strain that is unable to colonize the Mongolian gerbil (69).

Expression of pfr is tightly regulated by Fur. In keeping with

this finding, we detected strong induction of *fur* expression in the iron-starved cultures (Tables 1 and 2). *pfr* was strongly regulated only in the stationary phase, suggesting that iron-regulated expression of this gene is connected to the growth phase (Fig. 2). This view is consistent with the recent findings that the expression levels of *pfr* increase dramatically during the late exponential phase in a non-iron-chelated rich medium while the level of *fur* remains fairly constant (63). This is presumably because during exponential growth, iron is rapidly being utilized and thus sufficient quantities are not present for storage. In contrast, as the cells reach stationary phase and are no longer actively multiplying, iron levels become high enough to require storage of intracellular iron for later utilization during periods of iron starvation and to prevent toxicity (1).

(ii) Genes involved in translation. Analysis of the factors repressed by the removal of iron reveals that the majority of genes involved in translation were affected in exponential phase. Repression of many of these genes is most probably a direct result of the fact that the concentration of DPP used in this study resulted in cessation of growth in these cultures (data not shown). Since the bacteria are no longer dividing, the expression of factors that would be involved in growth was promptly down-regulated. This is evidenced by the fact that the largest class of repressed genes contains those predicted to play a role in translation. Included among these were genes encoding many ribosomal proteins and the translation initiation factor EF-1 (Table 1). It is perhaps for this reason that several genes involved in diverse forms of energy metabolism were repressed (Tables 1 and 2).

(iii) Genes encoding membrane proteins. Several genes that encode membrane proteins were also repressed in the exponential-phase culture. These include an inner membrane protein encoded by HP1450 and three outer membrane proteins (OMPs). The 64 identified OMPs of the two sequenced H. pylori strains have been recently divided into five paralogous families based on sequence similarity and predicted secondary structure (3). The major OMP family is composed of two subfamilies: the Hop and the Hor proteins. The Hop subfamily contains the largest number of OMPs, including several factors that are predicted to function as porins (21, 25) on adhesins (43). Two of the OMPs (HP0009 and HP0912;encoded by hopZ and hopC) repressed in exponential phase belong to the Hop subfamily, while another Hop protein (HP1177; encoded by omp27) was induced on iron starvation (Table 1). This opposing regulation perhaps suggests a functional shift in porin activity. Little is known about the selectivity of most of the Hop porins. Alterations in expression of the Hop proteins, specifically those acting as porins, would presumably affect the permeability of the outer membrane and may play a previously unappreciated role in iron uptake and/or storage.

(iv) Miscellaneous. Among the other five genes that were specifically repressed in the stationary phase are *dnaK*, which encodes the chaperone and heat shock protein 70 (Hsp70) and a conserved hypothetical protein, HP0388 (Table 2), that shows homology to predicted proteins in various pathogenic bacteria, such as *Haemophilus influenzae*, *Campylobacter jejeuni*, and *Vibrio cholerae*. Determination of the function of this hypothetical protein will be of interest in the future because its similar pattern of expression to *pfr* may suggest that it plays a

TABLE 1. Log-phase iron-regulated genes of H. pylori

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Gene designation ^a TIGR ASTRA		Gene function and name		Maximum fold change ^b in:	
				Expt 2	
Repressed genes					
Amino acid biosynthesis					
HP1229	JHP1150	Aspartokinase, <i>lysC</i>	0.4	0.4	
Cell envelope/Surface structures					
HP0009	JHP0007	Putative OMP <i>hopZ</i>	0.4	0.4	
HP0351	JHP0325	Flagellar basal-body M-ring protein, fliF	0.3	0.3	
HP0655	JHP0600	Putative OMP	0.4	0.4	
HP0858	JHP0792	ADP-heptose synthase, <i>rfaE</i>	0.4	0.5	
HP0870	JHP0804	Flagellar hook, flgE	0.4	0.5	
HP0912	JHP0848	OMP, hopC	0.4	0.5	
#HP1119	JHP1047	Flagellar hook-associated protein 1 (HAP1), flgK	0.3	0.4	
HP1275	JHP1196	Phosphomannomutase, algC	0.3	0.4	
HP1450	JHP1343	60-kDa inner membrane protein	0.4	0.5	
Cellular processes					
*HP0103	JHP0095	Methyl-accepting chemotaxis protein, <i>tlpB</i>	0.4	0.3	
HP0522	JHP0471	cag PAI protein, cag3	0.3	0.4	
HP1126	JHP1055	Colicin tolerance-like protein, <i>tolB</i>	0.2	0.4	
HP1300	JHP1220	Preprotein translocase subunit, secY	0.3	0.4	
HP1452	JHP1345	Thiophene and furan oxidizer, tdhF	0.3	0.5	
DNA metabolism-restriction/modification	1				
HP0481	JHP0433	Adenine-specific DNA methyltransferase, MFOKI	0.5	0.5	
Energy metabolism					
HP0509	JHP0459	Glycolate oxidase subunit, glcD	0.4	0.4	
HP0574	JHP0521	Galactosidase acetyltransferase, <i>lacA</i>	0.5	0.4	
HP1134	JHP1062	ATP synthase F1, subunit alpha energy metabolism, <i>atpA</i>	0.3	0.5	
Hymothetical					
Hypothetical HP0248	H1D0222	Conserved hymothetical protein	0.2	0.3	
HP0248 HP0270	JHP0233 JHP0255	Conserved hypothetical protein	0.2	0.3	
HP02/0 HP0466	JHP0418	Conserved hypothetical protein	0.3	0.4	
HP0554	JHP0501	Conserved hypothetical protein	0.3	0.4	
HP0659	JHP0604		0.4	0.3	
HP0783	JHP0720		0.4	0.4	
#HP0838	JHP0776		0.4	0.3	
HP0863	JHP0797		0.3	0.5	
HP1124	JHP1053		0.3	0.4	
HP1127	JHP1056		0.5	0.4	
HP1233	JHP1154		0.3	0.4	
Other categories, adaptations and					
atypical conditions					
HP1228	JHP1149	Invasion protein, putative dGTP pyrophosphohydrolase, invA	0.3	0.3	
Purine ribonucleotide biosynthesis					
HP0255	JHP0239	Adenylosuccinate synthetase, purA	0.5	0.4	
Tuescalistics					
Transcription #HP0550	JHP0497	Transcription termination factor, <i>rho</i>	0.2	0.3	
		•			
Translation	II ID0116	Pibosomal protoin I 20120	0.2	0.4	
HP0126 HP0247	JHP0116 JHP0232	Ribosomal protein L20, <i>rpl20</i> ATP-dependent RNA helicase, DEAD-box family, <i>deaD</i>	0.3 0.2	0.4 0.4	
	JHP0602	Putative processing protease			
#HP0657 HP0658	JHP0602 JHP0603	Glu-tRNA amidotransferase, subunit b, gatB	0.4 0.4	0.4 0.3	
HP0886	JHP0818	Cysteinyl-tRNA synthetase, cysS	0.4	0.5	
HP1298	JHP1218	Translation initiation factor EF-1, <i>infA</i>	0.4	0.5	
HP1299	JHP1219	Methionine aminopeptidase, map	0.3	0.5	
HP1304	JHP1224	Ribosomal protein L6, <i>rpl6</i>	0.5	0.5	
HP1305	JHP1225	Ribosomal protein S8, rps8	0.5	0.5	
HP1306	JHP1226	Ribosomal protein S14, rps14	0.4	0.5	
HP1312	JHP1232	Ribosomal protein L16, rpl16	0.4	0.5	
*HP1313	JHP1233	Ribosomal protein S3, rps3	0.3	0.4	
HP1314	JHP1234	Ribosomal protein L22, rpl22	0.3	0.4	
*HP1315	JHP1235	Ribosomal protein S19, rps19	0.4	0.4	
HP1554	JHP1445	Ribosomal protein S2, rps2	0.4	0.4	

TABLE 1—Continued

Gene designation ^a TIGR ASTRA		Gene function and name		Maximum fold change ^b in:	
				Expt	
Transport and binding proteins					
HP0497	JHP0449	Putative sodium- and chloride-dependent transporter	0.3	0.5	
HP1130	JHP1058	Biopolymer transport protein, <i>exbB</i>	0.3	0.3	
*HP1400	JHP1426	Iron(III) dicitrate transport protein, fecA3	0.3	0.5	
Induced genes					
Amino acid biosynthesis					
HP0134	JHP0122	3-Deoxy-D-arabino-heptulosonate-7-phosphate synthase, <i>aroF</i>	3.7	3.7	
#HP0106	JHP0098	Cystathionine gamma-synthase, <i>metB</i>	2.6	3.0	
#HP0107	JHP0099	Cysteine synthetase, <i>cysK</i>	2.9	2.6	
HP0626	JHP0570	Tetrahydrodipicolinate N-succinyltransferase, dapD	3.3	2.4	
*HP0649	JHP0594	Aspartate ammonia-lyase, aspA	2.1	2.5	
*#HP0672	JHP0615	Solute binding signature and mitochondrial signature protein, <i>aspB</i>	5.4	4.1	
Biosynthesis of cofactors, prosthetic					
groups, and carriers	TITDOSS:	Mall and I leave the last and the	2.5		
HP0577	JHP0524	Methylene tetrahydrofolate dehydrogenase, folD	2.7	2.4	
HP0665	JHP0610	Oxygen-independent coproporphyrinogen III oxidase, hemN	2.3	2.4	
HP0844 *#HP1582	JHP0782 JHP1489	Thiamine biosynthesis protein, <i>thi</i> Pyridoxal phosphate biosynthetic protein J, <i>pdxJ</i>	2.6 2.4	2.2 4.8	
	JHP1489	ryridoxai phosphate biosynthetic protein J, paxi	2.4	4.8	
Cell envelope and surface structures	HID0540	The colling A of A	2.1	2.1	
HP0601 HP0922	JHP0548 JHP0856	Flagellin A, flaA Toxin-like OMP, putative vacuolating cytotoxin (VacA)	3.1 3.6	2.1 3.0	
HP1052	JHP0373	paralog UDP-3-O-acyl-N-acetylglcosamine deacetylase, envA	3.2	2.5	
HP1177	JHP1103	OMP, omp27	2.4	2.3	
HP1192	JHP1117	Secreted protein involved in flagellar motility	2.7	2.6	
HP1456	JHP1349	Membrane-associated lipoprotein, <i>lpp20</i>	5.5	3.5	
HP1469	JHP1362	OMP, omp31	2.0	2.1	
*HP1501	JHP1394	OMP, omp32		2.9	
Cellular processes HP0243	JHP0228	Noutrophil activating protein (heatonicformitin) ugn 4	4.0	2.2	
HP0243 HP0541	JHP0228 JHP0489	Neutrophil-activating protein (bacterioferritin), napA cag PAI protein, cag20	3.6	3.2 2.1	
HP0547	JHP0495	cag PAI protein, cag20 (cagA)	3.5	2.5	
HP0875	JHP0809	Catalase, katA	2.6	2.3	
HP0979	JHP0913	Cell divison protein, GTPase in circumferential ring		2.3	
*HP1024	JHP0400	formation, <i>ftsZ</i> Cochaperone-curved DNA binding protein A, <i>cbpA</i>	4.7	2.2	
HP1069	JHP0356	Cell division protein, ftsH	4.6	3.0	
HP1431	JHP1322	16S rRNA (adenosine- N^6N^6)-dimethyltransferase, $ksgA$	2.4	2.7	
HP1563	JHP1471	Alkyl hydroperoxide reductase, tsaA		2.3	
Central intermediary metabolism					
HP0020	JHP0018	Carboxynorspermidine decarboxylase, nspC	2.2	2.3	
*HP0026	JHP0022	Citrate synthase, gltA	2.3 6.5	2.2	
*#HP0294	JHP0279	Aliphatic amidase energy metabolism, <i>amiE</i>		21.4	
HP0589 #HP0642	JHP0537 JHP0586	Ferredoxin oxidoreductase, alpha subunit, <i>oorA</i> NAD(P)H-flavin oxidoreductase	2.2 2.5	2.0 2.8	
HP0779	JHP0716	Aconitase B, acnB	4.9	2.4	
*HP0900	JHP0837	Hydrogenase expression/formation protein, <i>hypB</i>	3.2	4.1	
*HP0943	JHP0878	D-Amino acid dehydrogenase, <i>dadA</i>	3.2	2.7	
*HP1186	JHP1112	Carbonic anhydrase	2.3	2.2	
*HP1238	JHP1159	Aliphatic amidase, <i>amiF</i>	2.3	2.9	
HP1398	JHP1428	Alanine dehydrogenase, <i>ald</i>	3.9	2.8	
*HP1399	JHP1427 #JHP0585	Arginase energy metabolism, <i>rocF</i> Putative 3-hydroxyacid dehydrogenase	4.0 2.2	2.4 3.3	
Fatty acid and phospholipid metabolism		, , , ,			
*HP0690	JHP0638	Acetyl coenzyme A acetyltransferase (thiolase), fadA	2.7	2.3	
*HP0871	JHP0805	CDP-diglyceride hydrolase, <i>cdh</i>	8.2	6.9	
HP1045		Acetyl-CoA synthetase, acoE	4.5	3.6	
Hypothetical HP0018	JHP0016		2.5	3.0	
HP0080	JHP0074		3.5	3.3	
	3111 00 / T	Continue			

TABLE 1—Continued

Gene designation ^a TIGR ASTRA		Gene function and name	Maximum fold change ^b in:	
				Expt 2
HP0097	JHP0089		5.1	2.4
HP0105	JHP0097	Conserved hypothetical protein	3.2	2.9
#HP0204	JHP0190	71	2.2	2.0
HP0305	JHP0290		2.3	2.3
*HP0309	JHP0294	Conserved hypothetical protein	5.0	4.6
HP0310	JHP0295	Conserved hypothetical protein	6.7	4.1
#HP0311	JHP0296)	4.6	2.3
*HP0318	JHP0301	Conserved hypothetical protein	2.7	3.9
HP0367	JHP1014		2.1	2.5
*#HP0614	JHP0557		2.1	2.7
HP0627	JHP0571		2.3	2.3
HP0628	3111 03 / 1		2.6	2.5
HP0641	JHP0584		2.0	2.3
*HP0710	JHP0649	Conserved hypothetical protein	3.8	3.6
*HP0746	JHP0683	conserved hypothetical protein	2.4	2.5
HP0920	JHP0854	Conserved hypothetical integral membrane protein	3.0	2.7
*HP0944	JHP0879	Conserved hypothetical protein	3.0	3.7
HP1391	JHP1436	conserved hypothetical protein	6.8	2.8
HP1394	JHP1433	Conserved hypothetical protein	2.1	2.7
HP1454	JHP1347	Conserved hypothetical protein	2.1	2.7
HP1455	JHP1348		4.1	3.5
HP1457	JHP1350		3.9	3.3
HP1527	JHP1416		3.4	2.1
#HP1588	JHP1494	Conserved hypothetical protein	2.5	2.1
#111 1300	J111 1494	Conserved hypothetical protein	2.3	2.1
Regulatory functions				
HP0224	JHP0210	Peptide methionine sulfoxide reductase, <i>msrA</i>	3.2	2.6
HP0714	JHP0652	RNA polymerase sigma-54 factor, rpoN	2.3	2.1
*#HP1027	JHP0397	Ferric uptake regulation protein, fur	3.9	4.1
Translation				
HP0403	JHP0978	Phenylalanyl-tRNA synthetase, alpha subunit, pheS	2.0	2.5
HP0570	JHP0517	Aminopeptidase a/I, pepA	5.0	4.3
*HP1012	JHP0411	Putative zinc protease, pqqE	2.5	2.2
Transport and binding proteins				
*#HP0686	JHP0626	Iron(III) dicitrate transport protein, fecAI	2.1	3.3
HP0715	JHP0653	ABC transporter, ATP-binding protein	5.7	3.7
*#HP0807	JHP0743	Iron(III) dicitrate transport protein, fecA2	4.1	4.3
HP0940	JHP0875	Amino acid ABC transporter, periplasmic binding protein,	2.1	2.4
111 05 10	3111 0075	vckK	2.1	2
*HP1017	JHP0406	Amino acid permease, <i>rocE</i>	3.2	2.2
HP1091	JHP0334	Alpha-ketoglutarate permease, <i>kgtP</i>	2.8	2.5
HP1174	JHP1101	Glucose/galactose transporter, gluP	4.0	2.8
HP1491	JHP1384	Phosphate permease	3.6	2.4
Unknown function				
HP0390	JHP0991	Adhesin-thiol peroxidase, tagD	3.8	3.3
HP0569	JHP0516	GTP binding protein, gtp1	2.1	2.6
HP0625	JHP0569	Protein E, gcpE	3.3	3.4
	3111 0307		0.0	3.1

a*, these genes show differential regulation in both log- and stationary-phase cultures; #, these genes show differential regulation in the iron add-back experiment with the opposite change in expression.

role in iron sequestration. Extracellularly localized Hsp70 has been suggested to function as a sulfatide binding adhesion for *H. pylori* that facilitates attachment to sulfated glycolipids within the stomach mucus layer (28). Expression of *dnaK* is repressed by the HspR repressor protein (56), which we show here to be strongly induced by iron limitation, along with CbpA, encoded in the same operon (Table 2). We do not currently understand the significance of repression of *dnaK* on iron starvation; however, if surface-localized Hsp70 is in fact involved in mucosal adherence, decreased concentrations of iron may serve as a signal that promotes decreased adherence

and also promotes relocalization of *H. pylori* to a more suitable environment within the gastric mucosa. In support of this, we observed effects on genes involved in chemotaxis (*tlpB*) and components of the flagellar apparatus. These are discussed in greater detail below.

Iron starvation-induced genes. (i) Transport and binding. As expected, many genes that encode transport and binding proteins predicted to be involved in iron uptake were found to be regulated in our study. These include *fecA1*, *fecA2*, *fecA3*, and *frpB1*, which encode OMPs; *exbB*, which encodes a cytoplasmic membrane protein; and *ceuE1* and *ceuE2*, which en-

^b Due to space limitations, the maximum fold change is given for only one of the time points analyzed per experiment.

TABLE 2. Stationary-phase iron-regulated genes of H. pylori

Gene designation ^a		Gene function and name		ium fold ge ^b in:
TIGR	ASTRA			Expt 2
Repressed				
Amino acid biosynthesis #HP0652	JHP0597	Phosphoserine phosphatase, serB	0.2	0.2
#111 0032	3111 0377	Thosphosetine phosphatase, serb	0.2	0.2
Cellular processes, chaperones	IIID0101		0.4	0.5
HP0109	JHP0101	Chaperone and heat shock protein 70, dnaK	0.4	0.5
Energy metabolism				
#HP0632	JHP0575	Quinone-reactive Ni/Fe hydrogenase, large subunit, <i>hyaB</i>	0.4	0.4
#HP0633	JHP0576	Quinone-reactive Ni/Fe hydrogenase, cytochrome b subunit, hyaC	0.5	0.5
Hypothetical	****			
#HP0388	JHP0993	Conserved hypothetical protein	0.3	0.4
Transport and binding				
#HP0653	JHP0598	Nonheme iron-containing ferritin, pfr	0.2	0.1
T 1				
Induced Amino acid biosynthesis				
*HP0649	JHP0594	Aspartate ammonia-lyase, aspA	5.9	3.5
*#HP0672	JHP0615	Solute binding signature and mitochondrial signature protein, aspB	5.8	5.4
Biosynthesis of cofactors, prosthetic				
groups, and carriers			2.4	
HP0804	JHP0740	GTP cyclohydrolase II/3,4-dihydroxy-2-butanone 4-phosphate synthase, <i>ribAB</i>		2.0
HP1118	JHP1046	Gamma-glutamyltranspeptidase, ggt	3.3	2.5
*#HP1582	JHP1489	Pyridoxal phosphate biosynthetic protein J, pdxJ	8.6	6.6
#HP1583	JHP1490	Pyridoxal phosphate biosynthetic protein A, <i>pdxA</i>	6.4	3.9
Cell envelope and surface structures	11100105		4.0	2.1
HP0115 HP0175	JHP0107 JHP0161	Flagellin B, <i>flaB</i> Cell binding factor 2, putative peptidyl-prolyl <i>cis-trans</i> isomerase	4.0 2.4	2.1 2.7
*HP1501	JHP1394	OMP, <i>omp32</i>		2.1
Cellular processes				
*HP0103	JHP0095	Methyl-accepting chemotaxis protein, tlpB	2.0 2.6	4.9
HP0887	JHP0819	<i>U y y</i>		2.3
*HP1024	JHP0400	Cochaperone-curved DNA binding protein A, cbpA	4.6	3.5
Central intermediary metabolism	11100065		2.5	2.1
HP0072 HP0073	JHP0067 JHP0068	Urease beta subunit, <i>ureB</i> Urease alpha subunit, <i>ureA</i>	2.5 2.7	2.1 2.3
*HP0900	JHP0837	Hydrogenase expression/formation protein, <i>hypB</i>	2.6	2.1
*HP1186	JHP1112	Carbonic anhydrase	3.4	2.5
Energy metabolism				
*HP0026	JHP0022	Citrate synthase, gltA	2.6	2.2
HP0176	JHP0162	Fructose-bisphosphate aldolase energy metabolism, fba Ferredoxin energy metabolism	2.5 2.9	3.0
HP0277 *#HP0294	JHP0262 JHP0279	Aliphatic amidase, <i>amiE</i>	25.3	3.7 17.0
HP0691	JHP0637	3-Oxoadipate coenzyme A-transferase subunit A, <i>scoA</i>	3.6	3.7
HP0692	JHP0636	3-Oxoadipate coenzyme A-transferase subunit B, scoB	3.8	3.3
*HP0943 *HP1238	JHP0878 JHP1159	D-Amino acid dehydrogenase, <i>dadA</i> Aliphatic amidase, <i>amiF</i>	3.8 6.3	2.5 4.1
*HP1399	JHP1427	Arginase energy metabolism, rocF	2.6	2.1
HP1538	JHP1461	Ubiquinol cytochrome c oxidoreductase, cytochrome c_1 subunit, $petC$	2.2	2.1
HP1539	JHP1460	Ubiquinol cytochrome c oxidoreductase, cytochrome b subunit, $fbcH$	2.6	2.6
Fatty acid and phospholipid metabolism				
HP0416 *LIP0600	JHP0968	Cyclopropane fatty acid synthase, <i>cfa</i>	3.9	3.3
*HP0690 *HP0871	JHP0638 JHP0805	Acetyl coenzyme A acetyltransferase (thiolase), fadA CDP-diglyceride hydrolase, cdh	3.7 3.5	4.0 2.2

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TABLE 2—Continued

Gene designation ^a TIGR ASTRA		Gene function and name		Maximum fold change ^b in:	
				Expt	
Hypothetical					
*HP0309	JHP0294	Conserved hypothetical protein	3.3	2.6	
*HP0318	JHP0301	Conserved hypothetical protein	4.8	4.9	
HP0595	JHP0542		2.6	2.0	
*#HP0614	JHP0557		2.9	2.2	
HP0693	JHP0635	Conserved hypothetical integral membrane protein, atoE	3.2	2.8	
*HP0710	JHP0649	Conserved hypothetical protein	2.6	2.	
HP0719	JHP0657	•	2.1	3.:	
HP0721	JHP0658		2.3	2.	
*HP0746	JHP0683		2.1	2.	
HP0773	JHP0710		2.0	2.	
HP0914	JHP0850	Putative OMP	2.1	2.	
*HP0944	JHP0879	Conserved hypothetical protein	3.5	3.	
HP1173	JHP1100	conserved hypothetical protein	2.1	2.	
HP1182	JHP1108	Conserved hypothetical protein	2.6	2.:	
HP1507	JHP1400	Conserved hypothetical ATP binding protein	2.6	2.	
Other categories, adaptations, and atypical conditions					
HP1496	JHP1389	General stress protein, putative ribosomal protein L25, ctc	2.2	2.2	
Purines, pyrimidines, nucleosides, and nucleotides					
HP0680	JHP0621	Ribonucleoside-diphosphate reductase 1 alpha subunit, $nrdA$	2.3	2.2	
Regulatory functions					
HP0166	JHP0152	Response regulator, ompR	2.4	2.4	
HP1025	JHP0399	Putative heat shock protein, putative <i>hspR</i>		2.5	
*#HP1027	JHP0397	Ferric uptake regulation protein, fur		3.0	
Translation					
HP0177	JHP0163	Translation elongation factor EF-P, efp	2.1	2.:	
*HP1012	JHP0411	Protease, putative zinc protease, <i>pqqE</i>	2.1	2.:	
HP1019	JHP0405	Serine protease, htrA	2.1	4.	
*HP1313	JHP1233		2.3	2.	
*HP1315	JHP1235	Ribosomal protein S3, <i>rps3</i> Ribosomal protein S19, <i>rps19</i>	2.3	2.	
To a second and him him has a second since					
Transport and binding proteins	IIID0120	I Lagtata narmanca latD	2.7	2	
HP0140	JHP0128	L-Lactate permease, <i>lctP</i> Discretile APC transporter ATP his discrepation transport	2.7	2.0	
HP0301	JHP0286	Dipeptide ABC transporter, ATP-binding protein, dppD	2.5	2.	
*#HP0686	JHP0626	Iron(III) dicitrate transport protein, fecA1	6.4	4.0	
*#HP0807	JHP0743	Iron(III) dicitrate transport protein, fecA2	8.2	3.	
#HP0876	JHP0810	Iron-regulated OMP, frpB1	22.0 2.7	10.8	
HP0942	JHP0877	D-Alanine glycine permease, dagA		2.5	
*HP1017	JHP0406	Amino acid permease, <i>rocE</i>	2.4 2.6	2.4	
HP1172	JHP1099	glnH		3.1	
#HP1339	JHP1258	Biopolymer transport protein, <i>exbB</i>		2.5	
*HP1400	JHP1426			5.5	
#HP1432	JHP1321	Histidine- and glutamine-rich protein	5.4	2.4	
#HP1561	JHP1469	Iron(III) ABC transporter, periplasmic iron-binding protein, ceuE1	3.7	3.2	
HP1562	JHP1470	Iron(III) ABC transporter, periplasmic iron-binding protein, ceuE2	3.6	2.0	
Unknown function *HP1104	JHP1030	cinnamyl-alcohol dehydrogenase EL13-2, cad	3.1	3.2	

a*, these genes show differential regulation in both log- and stationary-phase cultures; #, these genes show differential regulation in the iron add-back experiment with the opposite change in expression.

b Due to space limitations, the maximum fold change is given for only one of the time points analyzed per experiment.

code periplasmic binding proteins (Tables 1 and 2). frpB1 was one of the most strongly regulated genes in the study, showing greater than 10-fold induction in the stationary-phase culture. Iron starvation affected the expression of several of these genes in a growth phase-dependent manner, since while both fecA1

and fecA2 were strongly induced in both stationary-phase and exponential-phase cultures, transcription of fecA3 showed differential regulation depending on the growth status of the cells. In the stationary phase, fecA3 was induced in a similar manner to its counterparts; however, in the exponential phase,

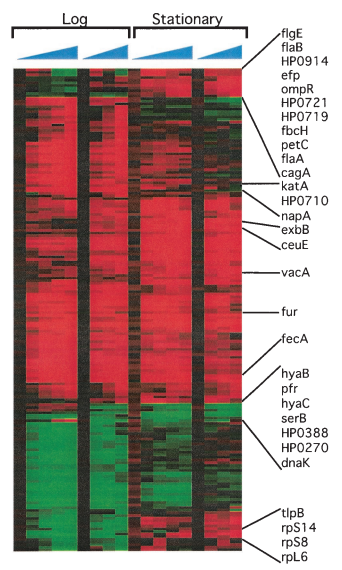


FIG. 2. Microarray analysis of the iron starvation response of *H. pylori*. A cluster diagram shows the expression profile of all 183 genes meeting the filter criteria (explained in Materials and Methods) after addition of the iron chelator. Two independent experiments for each growth phase are depicted, and relative expression patterns are shown for each time point. Increasing time postchelation is indicated by the blue triangles above the array data. Red indicates an increase in expression, while green indicates reduced expression. Representative genes are listed, and their relative location is indicated by lines. A complete list of the regulated factors and the maximum relative changes in expression can be found in Tables 1 and 2.

it was repressed by iron starvation. In addition, expression of ceuE1, ceuE2, and exbB is strongly regulated only in the stationary phase of growth. These data are in contrast with a recent report suggesting that expression of fecA3, ceuE1, and ceuE2 are unchanged by iron limitation (67). van Vliet et al. showed by spot blot hybridization that expression of these three genes remained virtually unchanged in H. pylori grown under iron-depleted conditions (67). Direct comparison of our results to theirs is confounded by the fact that while we analyzed both exponentially growing and stationary-phase cells, van Vliet et al. collected RNA from a single time point in the

growth of their strain and do not report which stage of growth this point represents. This observed discrepancy between the two studies may reflect growth phase differences in expression of the genes in question.

(ii) Virulence factors. Like other pathogenic bacteria, H. pylori seems to use iron limitation as a signal to regulate the expression of virulence factors. We observed induced expression of a number of genes that encode known virulence factors of H. pylori. These include cag20 and cag26 (cagA), both of which are located on the *cag* pathogenicity island (PAI). The PAI encodes a novel type IV secretion apparatus in some strains of H. pylori (2, 11), and it is well accepted that H. pylori strains carrying the PAI are more likely to be associated with serious manifestations of H. pylori infection (74). This is hypothesized to be because components of the island function in the delivery of CagA to the host cell, where it is inserted into the plasma membrane (52, 58) and induces striking morphological changes in the infected cell (51). Low concentrations of iron like those encountered within the host environment may serve as a signal that tells the bacterium that it is in a location where the production of virulence factors would be beneficial and may potentially induce the release of cellular components, such as iron, from damaged host cells.

Additionally, we observed increased expression of vacA, which encodes the vacuolating cytotoxin, VacA. VacA is a secreted protein that is internalized by host cells (38) and results in vacuolar degeneration that leads to the accumulation of large, nonfunctional endosomal-lysosomal hybrids (14, 35). The importance of VacA intoxication for the disease process remains elusive, but vacuolization of cells in human biopsy samples has been observed (10, 23). It was recently shown that vacA mutants have a 50% infectious dose that is 100-fold higher than that of an isogenic wild-type strain and show a significant colonization defect when coinfected with a wild-type strain (48). This suggests that VacA plays a role in establishing colonization, although the nature of this role is not immediately evident. Based on our and other (61) findings that vacA expression is up-regulated by iron starvation, one might propose that increased toxin production and the resulting increased cellular damage may lead to the release of host cell iron stores. Accordingly, it has been shown that administration of purified VacA to polarized T84 cell monolayers results in the increased permeability and release of Fe³⁺ and Ni²⁺ (45). Thus, toxin production and activity may facilitate the acquisition of iron by bacteria and promote bacterial survival in the host environment.

Induced expression of *napA*, which encodes the neutrophilactivating protein HP-NAP, was observed in the exponential-phase iron-depleted cultures. An independent study by Cooksley et al. recently showed that HP-NAP accumulation is influenced by Fur (13), once again confirming that our experimental design identified factors whose expression is affected by iron availability. HP-NAP was named for its ability to promote neutrophil adhesion to endothelial cells (24). It has been classified as a virulence factor and shown to be a major antigen in the human immune response to *H. pylori* (49). Additionally, is has been suggested to be a likely component for inclusion in an *H. pylori* vaccine due to its ability to provide colonization protection in mice orally immunized with purified HP-NAP (19, 49). Neutrophil activation and moderate levels of inflam-

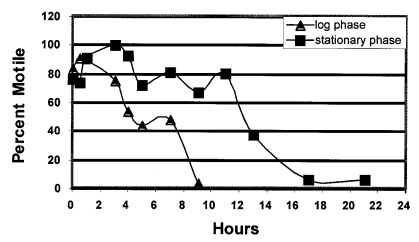


FIG. 3. Growth phase-dependent effect of iron starvation on *H. pylori* motility. Iron chelator was added to both exponential- and stationary-phase cultures of *H. pylori*, and the response of each culture was monitored by video microscopy as described in Materials and Methods. The percent motility of each culture over time is indicated.

mation induced by *H. pylori* are likely to play critical roles in bacterial colonization because they promote cellular damage that results in the leakage of nutrients from host cells (8). Thus, as with VacA, increased expression of HP-NAP on iron starvation might serve as a means of releasing iron from host cells. Further support for this is evident in the fact that the HP-NAP has strong in vitro bacterioferritin activity, binding up to 500 atoms of iron (64). Due to the important nature of this virulence factor and the predicted Fur binding site found by the bioinformatic analyses performed by us and by Cooksley et al. (13) (discussed below), future studies will attempt to discern the importance of this iron-dependent regulation.

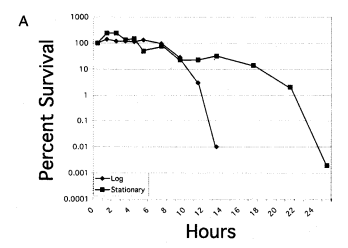
(iii) Nitrogen metabolism. The most highly regulated gene in the stationary-phase cultures was amiE, which encodes an aliphatic amidase. It, along with amiF, which encodes a formamidase, were up-regulated to various degrees by iron starvation, regardless of the growth status of the culture (Tables 1 and 2). As a group, bacterial amidases catalyze the hydrolysis of short-chain aliphatic amides to ammonia and the corresponding organic acid and play a valuable role in nitrogen metabolism (55). This is particularly true in H. pylori since ammonia not only is the preferred source of nitrogen for the synthesis of amino acids, pyrimidines, and purines but also contributes to epithelial cell damage and chemotactic motility and is a crucial component of the acid resistance pathway for the bacterium (30, 40, 50, 71). Previous characterization of the AmiE and AmiF amidases has indicated that they show marked substrate specificity and that their activity is dependent on the activities of other enzymes of the nitrogen metabolism pathways of H. pylori, namely, those of urease and arginase (55). Expression of amiF and amiE strongly correlates with growth phase-dependent patterns of expression of fecA and frpB, perhaps suggesting that they have similar regulatory signals (63). Recently, van Vliet et al. reported that Fur regulates the transcription, expression, and activity of AmiE and indirectly regulates the activity of AmiF (66). Therefore, our finding of strong iron-dependent regulation is not surprising. Perhaps what is surprising, though, is the apparently strong level of coregulation of genes involved in nitrogen metabolism and iron starvation. We found that in addition to amiE and amiF,

multiple genes that are involved in pathways of ammonia and urea production were iron regulated. These include genes encoding components of the urease enzyme, *ureA* and *ureB*; genes involved in arginine metabolism, *rocE* and *rocF*; and a component of the glutamine transporter, *glnH* (Tables 1 and 2). The reason for the coregulation of these two apparently very different metabolic pathways is currently unclear, but since ammonia causes epithelial cell damage, increased ammonia production may also serve as a mechanism by which *H. pylori* increases cellular damage and thus triggers the release of host cell iron stores. It is also likely that iron chelation affects iron-cofactored enzymes that play a role in nitrogen metabolism, and this might account to some degree for the apparent coregulation.

(iv) Flagellar genes and motility. Observation of the growth phase-dependent expression of components of the flagellar apparatus included increased expression of *flaA* in exponential-phase cells but increased *flaB* expression in stationary-phase cells (Tables 1 and 2). Previous analysis of the *H. pylori* flagellar filament has shown that it consists of a copolymer of FlaA and FlaB (27, 36, 60), where FlaA is the predominant subtype and FlaB is present as a minor constituent (33). Both FlaB and FlaA are necessary for full motility (32). In comparison to most other bacteria, the existence of two distinct flagellin subunits present in different amounts is somewhat unusual. It has been suggested that *H. pylori* has the ability to alter the relative levels of FlaA and FlaB in response to different envi-

TABLE 3. Comparison of the progression of motility during iron chelation compared to normal growth at both exponential and stationary phases

Exp	onential pha	se	Sta	tionary phas	e
Condition	Growth time (h)	% Motile	Condition	Growth time (h)	% Motile
Control	Start 12	82 66	Control	Start 12–13	66 34
Chelation	Start 9	87 4	Chelation	Start 12–13	76 38



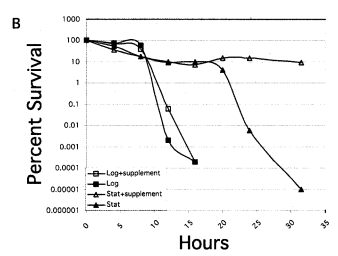


FIG. 4. (A) Iron starvation effects on cell viability. The viability of iron-depleted $H.\ pylori$ cells was monitored by plating for CFU. Iron chelator was added to both exponential- and stationary-phase cultures, and titers were taken periodically thereafter. The percent survival was calculated based on initial titers taken at the t=0 time point, and times indicated on the x axis indicate hours postchelation. (B) Effect of supplementation on cell viability. To determine whether decreased viability of iron-starved cells was due to expended stores of iron or of another nutrient, supplementation experiments were conducted. Both exponential- and stationary-phase cultures were maintained without supplementation as controls and showed kill curves similar to those depicted in panel A. For supplemented cultures, approximately 20% of the culture was removed every 4 h and replaced with previously chelated BB as described in Materials and Methods.

ronmental stimuli (57, 59) and two recent independent studies have demonstrated that transcription of *flaA* and *flaB* was strongly regulated in a growth phase-dependent manner (41, 63). Interestingly, in those studies *flaA* showed maximal expression in the stationary phase while *flaB* expression peaked early in the exponential phase. This is opposite to the effect we observed when we starved the cells for iron, thus suggesting that in addition to growth phase dependency, further levels of flagellar regulation are governed by environmental signals. By altering relative concentrations of the two flagellar subunits, the bacterium could potentially adapt to microenvironments by alteration of the filament ultrastructure. This may alter flagel-

lar stiffness or flexibility to better suit the particular environment being encountered.

To determine whether iron starvation of H. pylori showed a differential effect on motility depending on the growth phase of the culture, we conducted video microscopy of H. pylori cells exposed to iron starvation conditions for extended periods as described in Materials and Methods. Analysis of these bacterial populations showed that prior to chelation, roughly equivalent numbers of bacteria (approximately 75%) were motile in the exponential and stationary phase cultures (Fig. 3). On iron starvation, the exponential phase decreased in the number of motile cells. By 9 h, the entire culture was nonmotile. This is in contrast to a normal growth curve, where motility decreases much more slowly, remaining at approximately 60% even after 12 h of growth (Table 3). Morphologically, at the later time points the iron-restricted bacteria in the exponential-phase culture appeared coccoid (data not shown). Analysis of the stationary-phase culture revealed that virtually the entire culture became motile by 3 h postchelation, and that overall levels of motility were maintained for much longer periods. This is similar to what is observed in a normal growth curve (Table 3). Of note, more than 60% of the iron-restricted culture was still highly motile after 9 h when all exponential-phase bacteria appeared coccoid. The motility of the stationary-phase cells remained relatively high for approximately 14 to 16 h after iron starvation; moreover, visually these cells appeared helical and noncoccoid for extended periods (data not shown). These data indicate that iron starvation shows growth phase-dependent effects on motility in H. pylori. The significance of this in the in vivo environment is not immediately clear, but the ability to sense the surrounding environment and alter motility probably serves as a critical component of successful colonization because it would direct H. pylori to a suitable environment for colonization. Of note, a single chemotaxis gene (tlpB) was shown to be differentially regulated by iron starvation, perhaps suggesting a role for this gene in responding to iron concentration.

Growth phase effects on iron starvation susceptibility. Based on our observation that iron-starved exponential-phase cells become nonmotile and coccoid considerably earlier than stationary-phase cells, we hypothesized that the growth phase might play a role in susceptibility of *H. pylori* to iron starvation. To investigate this, we conducted a chelation time course experiment and calculated the percent survival of iron-starved exponential- and stationary-phase cells. As shown in Fig. 4A, both exponential- and stationary-phase cells displayed similar survival profiles for the first 8-10 h of the experiment. After this point, survival rapidly decreased in the exponential-phase cells. However, a significant proportion of the stationary-phase cells remained viable for more than 20 h, suggesting that intracellular iron stores of the stationary-phase bacteria were sufficient to support extended periods of iron depletion. To ensure that decreased viability of the bacterial cultures was due to lack of iron and not due to deficiency of some other nutrient, we supplemented both exponential- and stationary-phase ironstarved cultures with freshly chelated medium as described in Materials and Methods. To our surprise, while this had no effect on the viability profile of the exponential-phase cells, it completely abrogated the decline in viability that we had originally observed in the later time points of the stationary-phase

В ----AATAAT--T-A-consensus HP0073 TTTTAAATAATAATTAGT ureA HP0243 f napA GATTAATATATTAAAT HP0294 f amiE CCCACAATAATCATAATG HP0309 f hypothetical TGAATAATAGTAAGT HP0390 tagD GTTAGAATAATAGTCAAT HP0686 f fecA1 AAACTAATAATGGTTATT ribAB HP0804 f AAAGTAATAATCGTTATT HP0804 TATTCAATAATATTGAGT ribAB fecA2 HP0807 ATAAAAATAATTATTAAA HP0871 f cdh ATTAAAATAATCATAATG HP0876 f GTTTTAATAATAATTATT frpB1 HP0876 **AGTATAATAATTATTATT** frpB1 HP0912 omp20/hopC TTGTAAATAATGTTCAGT HP0920 hypothetical IMP CAATAATAATTTTAATT HP0921 f CAATAAATAATTTTAATT gap HP1027 f ATTAAAATAATAATGAGT fur gluP HP1174 f CTTAAAATAATTGTAATA gluP HP1174 GATATAATAATGATA amiF HP1238 f CTCGCAATAATTGTTATT exbB HP1339 AAGTCAATAATGATTAAG HP1527 hypothetical TAATTAATAATAATTAGT

FIG. 5. (A) Identification of a Fur binding site consensus sequence. Sequence alignment of the high-affinity Fur binding sites of the frpB (designated box 1 in reference 16) and fur (designated box I in reference 18) promoters, as well as the FURTA-Hp positive fecA2 promoter (27). Conserved nucleotides are indicated in bold and denoted by asterisks. (B) Putative identification of Fur boxes in predicted promoter regions of iron-regulated genes. Sequence alignment of all genes shows a perfect match to the determined consensus sequence. Genes are listed in chromosomal order, and HP designations represent those given by TIGR. The "f" or "r" following the HP number indicates the forward and reverse strand, respectively, on which the consensus match was found. Conserved nucleotides are indicated in bold.

culture; we observed a 6-log-unit difference in the number of viable cells at the latest time point at which measurements were made (Fig. 4B). This suggests that stationary-phase cells do not die due to a lack of iron but die because of a lack of some other unidentified nutrient and indicates that stationary-phase *H. pylori* cultures have a previously unappreciated tremendous capacity for survival when faced with iron starvation, possibly due to the ability of Pfr to accumulate large intracellular iron stores. This ability to survive lengthy periods of iron starvation could be of significant clinical importance and probably plays a role in the environment of the human stomach, where available iron is tightly regulated through sequestration by the host.

Bioinformatic analysis reveals putative Fur-regulated genes. Since Fur regulation is a key component of iron acquisition and sequestration and since very few genes have actually been proven to be directly regulated by Fur binding in *H. pylori*, we

performed a preliminary bioinformatic analysis of our differentially regulated genes to identify putative Fur binding sites in the predicted promoter regions. Using the sequence of the high-affinity Fur boxes from frpB, fur, and the FURTA-Hppositive fecA2 promoter (16, 17, 26), a candidate Fur box consensus sequence (AATAATNNTNA) for H. pylori was generated (Fig. 5A). Homology searches of the predicted promoter regions of the 183 differential regulated genes identified 18 genes that contained a perfect match to the consensus sequence (Fig. 5B). Of these, three genes contained more than one putative Fur box: gluP, ribAB, and frpB1. Analysis of the list revealed a strong enrichment for genes that are predicted to play a role in iron acquisition or have previously been proposed to be Fur regulated, thus suggesting that our bioinformatic approach was able to identify at least a subset of Fur-regulated genes. Included among these are fecA1, fecA2, exbB, ribAB, napA, and amiE. Also included in this list are

several genes whose putative regulation by Fur had not previously been hypothesized. Among these are several genes encoding hypothetical proteins, proteins of unknown function, and predicted membrane proteins. A goal of future research will be to directly investigate the Fur regulation of these genes.

Iron add-back. Finally, since the removal of iron from the exponentially growing culture of H. pylori results in growth cessation and thus has both direct and indirect effects on gene expression, we wished to conduct an additional approach to complement our growth phase dependence study. We therefore conducted microarray analysis of H. pylori grown in an iron-limited environment (see Materials and Methods) to examine the transcriptional effect of the addition of excess iron to these cells. We hypothesized that genes that showed altered expression might represent those that are the most strongly iron regulated. This iron add-back study revealed that 28 genes overlapped with the growth phase-dependent identified factors from the chelation experiments (denoted in Tables 1 and 2 by #). The 28 genes were equally distributed between the exponential- and stationary-phase lists of affected factors. Interestingly, multiple genes that have been previously shown to be iron regulated, such as vacA and napA (13, 61), were not identified by this method. This highlights the importance of using complementary conditions for comprehensive identification of regulons affected by a stimulus of interest.

Concluding remarks. A formidable challenge in understanding how bacterial pathogens colonize and cause disease in their hosts is the development of a more thorough understanding of the adaptive mechanisms used by these pathogens within the host. The gastric pathogen H. pylori undoubtedly encounters a number of environmental stresses within the human stomach where it colonizes. One of these stresses is likely to involve periods of iron starvation because the human body tightly sequesters available iron to prevent oxidative damage and to inhibit bacterial growth. In an effort to more thoroughly define the response of H. pylori to iron starvation, we used DNA microarrays to investigate global temporal changes in gene expression of both exponentially growing and stationary-phase cultures. We chose to investigate both growth phases as a result of a recent study suggesting that many of the genes of H. pylori that are differentially regulated as the bacteria progress through the growth phase are predicted to be iron regulated or to play some role in iron acquisition (63). We therefore hypothesized that iron starvation might have drastically different effects on the transcriptome of *H. pylori* depending on the growth phase of the bacteria. This was indeed the case, since we noted that many regulated genes showed growth phase-dependent changes in transcription. This result highlights the importance of considering growth phase when interpreting the results of future studies directed at understanding the transcriptional responses of *H. pylori*.

Future experiments to identify the role of each of the identified factors in adaptation to iron starvation should provide valuable information concerning the mechanisms used by *H. pylori* during colonization within the gastric environment of the human host. Since the ability to acquire and store iron is a crucial part of the ability of *H. pylori* to survive in its host and cause disease, investigation of some of these factors may facilitate the elucidation of novel targets for vaccine development and antimicrobial therapy.

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D. Scott Merrell and Lucinda J. Thompson contributed equally to this work.

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